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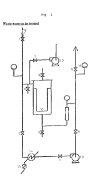
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(54) METHOD OF TREATING WASTEWATER CONTAINING HARDLY DECOMPOSABLE HARMFUL SUBSTANCES

(57) A method of treating hardly decomposable harmful substance comprising (A) a step for adding coaguiant into wastewater containing a hardly decomposable harmful substance. (B) a step for performing soliditiquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid, and (C) a step for decomposing the hardly decomposable harmful substance by bringing percuide into contact with the separated solid. Efficiently and seldy treating the hardly decomposable harmful substances by means of low cost facilities realizes.



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#### Description

#### TECHNICAL FIELD

5 (0001) The present invention relates to a method of treating wastewater containing hardly decomposable harmful substances. More particularly, the present invention relates to a method for treating wastewater which coagulates and as exparates hardly decomposable harmful substances from wastewater containing the hardly decomposable harmful substances, and efficiently decomposable harmful substances.

#### 10 BACKGROUND ART

[0002] Hardly decomposable substances such as dioxins known as the substance harmful to human body, i.e., hardly decomposable harmful substances are discharged into natural environment from incinerator for urban waste or industrial waste and various combustion facilities or machineries.

[0033] Moreover, various kinds of organic compounds in manufacturing process of chemical substance which will unintentionally give bad influence upon environment are discharged causing big social problems. Although the paeneation mechanism of chloridized aromatic compound such as dioxins, chlorophenol, chlorobenzene is not appearent, it is said that they generate in a treating process for wasting gas of low temperature in the presence of nonflammable carbon, air, molsture, inorpanic chlorine, etc.

0 [004] As sources of releasing wastewater containing these compounds, chloride base bleach facilities in Kraft pulp production plant, decomposition tacilities of wasted PCB or processed PCB, cleaning equipment for PCB contaminant or PCB processed product, cleaning equipment modifying fusion furnace to provide for manufacture of aluminum and aluminum alloy, wet type dust collection facilities, waste pit discharging sewage, etc., are known. Namely, a process using compounds containing choldreds has probability of generating the foreoing wastewater.

28 [0005] Further, standard of aqueous environment pollutant was revised by Ministry of Environment and as a result, organic compounds such as trichloroethylene, tetrachloroethylene, PCB were newly added to object substance of the environmental quality standard in which heavy metals were main entity until then. It is probable that a slight amount of harmful substance such as organophosphorus compound, PCB, trichloroethylene, dioxines, bisphenois and so on is contained in industrial wastewater, in addition to oils, chemical oxygen demand (COD), biochemical oxygen demand (BOD) and suspended substance (SS).

[0006] Conventionally, technologies for decomposing hardly decomposable harmful substances in wastewater which removes the hardly decomposable harmful substances as much as possible from the object wastewater containing the hardly decomposable harmful substances by means of a filtration equipment, a membrane separation method, etc., have been developed (refer to, for example, Japanese Unexamined Patent Application Laid-Open No. Hei 11-93395).

[0007] For the purpose of treating wastewater containing the hardly decomposable harmful substances as mentioned above, a filtration treatment, a bio-organic treatment or so is carried out as a pretreatment and then, an ozone treatment, an ultraviole irradiation treatment, a catalyst treatment, an activated carbon treatment or so is carried out as an aftertreatment. As thus described, it was necessary to decompose and remove the hardly decomposable harmful substances spending orgal labors and materials in the post.

[0008] Further, giving the ultraviolet irradiation treatment in example, it is a technology which can take advantage only about the reaction system in which the ultraviolet ray can transmit, and there is a problem that it cannot take advantage about solid or a liquid containing solid. Furthermore, it is necessary for the hardly decomposable harmful substances removed by the pretreatment to be processed specially into harmless in order to prevent the secondary contamination.

[0009] Accordingly, development of a decomposition treatment technology for these hardly decomposable harmful substances efficiently and without contaminating human body or surrounding environment again is eagerly desired.

## DISCLOSURE OF THE INVENTION

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[0010] An object of the present invention is to provide a method of treating wastewater containing hardly decomposable harmfull substances which coagulates and separates the hardly decomposable harmful substances from wastewater and efficiently decomposes the separated hardly decomposable harmful substances in the solid condition.

[0011] As a result of intensive extensive research and investigation made by the present inventors in order for echieving the object, it has been found that a solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid with a use of a coegulant, and decomposing the hardly decomposable harmful substance by bringing peroxide into contact with the separated solid achieve the object. Such being the case, the present invention has been accomplished on the basis of the foregoing lindings and information.

#### [0012] Namely, the present invention provides:

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- (1) A method of treating hardly decomposable harmful substance comprising:
  - (A) a step for adding coagulant into wastewater containing a hardly decomposable harmful substance, (B) a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid, and
  - (C) a step for decomposing the hardly decomposable harmful substance by bringing peroxide into contact with the separated solid:
- (2) The method of treating hardly decomposable harmful substance according to the above item (1), wherein the hardly decomposable harmful substance is at least one kind selected from the group consisting of dibenzodoxin haildes, dibenzoturan haildes, polychlorinated biphenyls, benzene haildes, alkylphenols, phenol haildes, alkane haildes. Alkene haildes, bithtalic esters, bischenols and polycyclic aromatic hydrocarbons:
- (3) The method of treating hardly decomposable harmful substance according to the above item (1), wherein the coagulant employed in step
  - (A) is at least one kind selected from the group consisting of polyacrylic acid, polyacrylamide resin, ferrous coagulant and aluminum-based coagulant;
- (4) The method of treating hardly decomposable harmful substance according to the above item (3), wherein the ferrous coagulant is at least one kind selected from the group consisting of ferric sulfate, polyferric cholde; chloride and polyferric chloride;
- (5) The method of treating hardly decomposable harmful substance according to the above item (3), wherein the aluminum-based coadulant is at least one of aluminum chloride and polyaluminum chloride:
- (6) The method of treating hardly decomposable harmful substance according to the above item (1), further adding coagulation promoter in the step (A):
- (7) The method of treating hardly decomposable harmful substance according to the above item (6), wherein the coagulation promoter is an inorganic porous material or an organic porous material;
- (8) The method of treating hardly decomposable harmful substance according to the above item (6), wherein the coagulation promoter is at least one kind selected from the group consisting of ion exchange resin, diatom earth, activated clay, zeolite, pearlite and activated carbon;
- (9) The method of treating hardly decomposable harmful substance according to the above item (1), wherein a filtration method, a centifugal separation method or a membrane separation method is employed in the step (8); (10) The method of treating hardly decomposable harmful substance according to the above item (1), wherein the peroxidic used in the step (6) is an oxidizing agent;
- (11) The method of treating hardly decomposable harmful substance according to the above item (10), wherein the oxidizing agent is at least one of persulfate and permanganate:
- (12) The method of treating hardly decomposable harmful substance according to the above item (11), wherein the persulfate is at least one kind selected from the group consisting of ammonium persulfate, sodium persulfate and potassium persulfate;
  - (13) The method of treating hardly decomposable harmful substance according to the above item (11), wherein the permanganate is at least one of sodium permanganate and potassium permanganate:
- (14) The method of treating hardly decomposable harmful substance according to the above item (1), wherein the peroxide used in the step (C) is at least one selected from the group consisting of salt of peroxo acid, hyperoxide and organic peroxide;
  - (15) The method of treating hardly decomposable harmful substance according to the above item (14), wherein the salt of peroxo acid is perborate or organic peroxide;
  - (16) The method of treating hardly decomposable harmful substance according to the above item (14), wherein the hyperoxide is at least one selected from the group consisting of potassium hyperoxide, sodium hyperoxide, barlum hyperoxide and magnesium hyperoxide;
    - (17) The method of treating hardly decomposable harmful substance according to the above item (14), wherein the organic peroxide is ditertiarybutylperoxide or dicumyl peroxide;
    - (18) A method of treating hardly decomposable harmful substance comprising:
      - (a) a step for adding an adsorbent into wastewater containing a hardly decomposable harmful substance, (B) a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid, and

- (C) a step for decomposing the hardly decomposable harmful substance by bringing peroxide into contact with the separated solid:
- (19) The method of treating hardly decomposable harmful substance according to the above item (18), wherein the adsorbent is at least one selected from the group consisting of ion exchange resin, diatom earth, activated clay zeolite, certific litania and activated carbon:
  - (20) A method of treating a hardly decomposable substance wherein the hardly decomposable substance adsorbed to a solid is brought into contact with peroxides in an amount of 100 moles or more per the hardly decomposable substance without conducting any desoration operation:
- (21) The method of treating a hardly decomposable substance according to the above item (20), wherein the solid is any one of soil, sludge, incinerated ash or adsorbent;
  - (22) The method of treating hardly decomposable harmful substance according to the above item (20), wherein in he hardly decomposable substance is at least one selected from the group consisting of dilberacodioxin halides, diberacofuran halides, polychlorinated biphenyls, benzene halides, alklylhenols, phenol halides, alkane halides
  - (23) The method of treating hardly decomposable harmful substance according to the above item (20), wherein the peroxide is at least one selected from the group consisting of salt of peroxo acid, hyperoxide and organic peroxide.
- (24) The method of treating hardly decomposable harmful substance according to the above item (23), wherein the salt of person acid is at least one selected from the group consisting of persulfate, permanganate, perborate and organic peroxide:
  - (25) The method of treating hardly decomposable harmful substance according to the above item (24), wherein the persulfate is at least one selected from the group consisting of ammonium persulfate, sodium persulfate and
- potassium persulfate;

  25 (26) The method of treating hardly decomposable harmful substance according to the above item (24), wherein
  the permanganate is any one of sodium permanganate and potassium permanganate;
  - (27) The method of treating hardly decomposable harmful substance according to the above item (23), wherein the hyperoxide is at least one selected from the group consisting of potassium hyperoxide, sodium hyperoxide, barfum hyperoxide and magnesium hyperoxide; and
- 30 (28) The method of treating hardly decomposable harmful substance according to the above item (23), wherein the organic peroxide is ditertiary butylperoxide or dicumyl peroxide.

#### BRIEF DESCRIPTIONOF THE DRAWINGS

- 35 [0013] Fig. 1 illustrates a block diagram of a solid-liquid separator with a bag filter employed in Examples and Comparative Examples.
  - [0014] In Fig. 1, reference numerals 1 to 9 indicate valves; reference numeral 10 indicates a bag filter; reference numeral 11 indicates a pressure gauge for measuring differential pressure of the filter; reference numerals 12 and 13 indicate pumps; reference numeral 15 indicates a temperature controlling trag; and reference numeral 16 indicates a drain line for the wastewater.

#### THE PREFERRED EMBODIMENT TO CARRY OUT THE INVENTION

- [0015] To begin with, the first aspect of the present invention provides a method of treating hardly decomposable 45 harmful substance comprising (A) a step for adding coagulant into wastewater containing a hardly decomposable harmful substance; (B) a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid; and (C) a step for decomposing the hardly decomposable harmful substance by bringing percoids into contact with the separated solid.
- [0016] Examples of the hardly decomposable harmful substances in the method of treating wastewater containing a hardly decomposable harmful substances (it may be described merely as "a method for treating wastewater" below) in the present invention include dibenzodioxin halides (dioxins), dibenzoduran halides, polychiorinated byhensyls (PCB), benzone halides, alikylphenols, phenol halides, alikene halides, alikene halides, phihalic esters, bischenols and polycyclic aromatic hydrocarbons.
- [0017] Typical examples of the dibenzodioxin halides include compounds such as 2,3,7,8-tetrachlorodibenzo-p-di-sion, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,6,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,5,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,8,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,8,7,8-pentachlorod-benzo-p-dioxin, 1,2,3,4,5,7,8-penta
  - [0018] Typical examples of the dibenzofuran halides include compounds such as 2,3,7,8-tetrachlorodibenzofuran, 1,2,3,7,8-pentachlorodibenzofuran, 2,3,4,7,8-pentachlorodibenzofuran, 1,2,3,4,7,8-hexachlorodibenzofuran,

- 1,2,3,6,7,8-hexachlorodibenzofuran, 1,2,3,7,8,9-hexachlorodibenzofuran, 2,3,4,6,7,8-hexachlorodibenzofuran, 1,2,3,4,6,7,8-hexachlorodibenzofuran, 1,2,3,4,6,7,8-hexachlorodibenzofuran, etc.
- [0019] As the polychlorinated biphenyls, there are Coplanar PCBs whose chlorine atom is substituted except at ortho position, specific examples include compounds such as 3,3',4,4'-fetrachloro biphenol, 3,3',4,4'-fe-pentachlorobiphenol, 3,3',4,4'-fe-pentachlorobiphenol, etc.
- [0020] Typical examples of the benzene halides include compounds such as chlorobenzene, dichlorobenzene, trichlorobenzene, tetra chlorobenzene, pentachlorobenzene, haxachlorobenzene, etc.
- [0021] Typical examples of the alkylphenosi include compounds such as t-bulylphenol, nonylphenol, octyl phenol, pentyl phenol, etc., and typical examples of the phenol halides include compounds such as chlorophenol, dichlorophenol, trichlorophenol, tertachlorophenol, pentachlorophenol, etc.
- [0022] "Typical examples of the alkane halides or the alkene halides include compounds such as dichloropropane, trichloroethane, trichloroethylene, tetrachloroethylene, dichloroethylene, etc., and examples of the phthalic esters include compounds such as dibutyl phthalate, butyl borzyl phthalate, di-2-ethylknyphthalate, and the phthalic esters include compounds such as dibutyl phthalate, butyl borzyl phthalate, di-2-ethylknyphthalate, and the phthalic esters include compounds such as dibutyl phthalate, butyl borzyl phthalate, di-2-ethylknyphthalate, and the phthalic esters include compounds such as dibutyl phthalate, but the phthalate of a compound such as dibutyl phthalate, but t
- [0023] Typical examples of the bisphenols include compounds such as 2,2-bis (4-hydroxyphenyl) propane (bisphenol A), 1,1-bis (4-hydroxyphenyl) cyclohexane, or so and typical examples of the heterocyclic aromatic hydrocarbons include benzopyrene, chrysene, benz anthracene, benzofluorenthene, picene, etc.
- [0024] The melihod for treating wastewater of the present invention is preferably applied to the hardly decomposable harmful substances such as halogenated dibenzodioxinas (dioxinas), polychlorinated biphonyts (PCB), benzene halides, slixyhonistides, biphonistic and dibenzodioxina halides.
- 20 [0025] In the present invention it is desirable to preparedly adjust pH of the wastewater containing the hardly decomposable harmful substances within the range of from 6 to 12 before coagulating the hardly decomposable harmful substances by adding the coagulant to the wastewater as the step (A). When pH is lower than 6, pipes of processing apparatus may corrode and when pH exceeds 12, neutralization of the wastewater at the last step will become troublesome. Detailed explanation about a basic substance for adjusting pH will be described below in a term about the basic substance.
  - [0026] Carrying out the step (A), that is, adding the coagulant into the pH adjusted wastewater after completing the above pH adjustment will coagulate the hardly decomposable harmful substances.
  - [0027] Regarding with the coagulant employed for the present invention, ferrous-based and/or aluminum-based inorganic ocaquinal is preferable. Typical examples of the ferrous-based coagulant include ferric sudfate, polyferric suifate, ferric chloride, polyferric chloride, etc., and typical examples of the aluminum-based coagulant include aluminum chloride, oby aluminum chloride, etc.
- [0028] Further, organic coagulant such as polyacrylic acid, polyacrylamide-based polymer and so on may be also employed. The foregoing coagulant may be employed alone or in combination of two or more kinds thereof. The addition amount of the coagulant is not particularly restricted and it may be appropriately settled depending on the kind of the coagulant is not particularly restricted and it may be appropriately settled depending on the kind of the coagulant employed or the coagulation component amount in the wastewater, however, it is generally settled within the range of from 0.0001 to 10 % by weight to the amount of the wastewater, preferably within the range of from 0.0001 to 1 % by weight.
- [0029] In the present invention, a coagulation promoter may be optionally employed in combination with the coagulant. Specific examples of the coagulation promoter employed in the present invention include an inorganic power.

  material such as diatom earth, zeolite, pearlife, activated clay and so on, or an organic porous material such as activated carbon, ion exchange resin, etc. The kind of the coagulation promoter may be selected depending on the wastewater to be treated. They may be used alone or in combination of two or more kinds thereof. The addition amount of the coagulation promoter is not particularly restricted and is appropriately settled depending on the kind of the coagulation promoter employed or on the amount of coagulation component in the wastewater.
- 45 [0030] As a method for adding the coagulation promoter, there are a precoat method forming a cake any fiftering. In the precoat method forming a cake to fiftering, in the precoat method, the addition amount may be decided from filtration area, and in the body feed method, the addition amount may be decided from filtration area, and in the body feed method, the addition amount may be decided making a turbidity of wastewater in a rule of thumb.
  - [0031] The step (B) in the method for treating wastewater is a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substances from a liquid.
- [0032] In the solid-liquid separation step, for example, a membrane separation method, a filtration method, a centrifugal separation method and so on are employable as the solid-liquid separation method. In the membrane separation method, various kinds of membrane separation method, various kinds of membrane are microfiltration (MP) membrane, an ultrafiltration (MP) membrane, a morse osmosis (RO) membrane, and so on are usable as the separation membrane. So Among these, MF membrane and UF membrane are preferable because they are efficient even under low pressure and they are economic in the viewpoint of a cost.
  - [0033] As a material for MF membrane, each resin such as cellulose-base, polyamide-base, polysulfone-base, polyelene-base, cellulose acetate-base, polyelher sulfone-base, polyacrylonitrile-base and so on may be used.

[0034] Further, as a material for UF membrane, each resin such as polyamide-base, polysulfone-base, polypropyiene-base, polyvinylpymolidone-base and so on may be also used similarly as the material for MF membrane. In addition to the above various membranes, a bag filter is used as preferred means.

[0035] On the other hand, a sand filtration method may be employable as filtration method, and a cross flow system may be employable as an entire filtration system.

[0036] The step (C) in the method for treating wastewater in accordance with the present invention is a step for decomposing the hardly decomposable harmful substances by bringing peroxide into contact with the separated sold. In the method for treating wastewater in accordance with the present invention, the hardly decomposable harmful substances coagulated into the above solid are able to be oxidatively decomposed in the state of solld by peroxide without carrying out desorption operation. Accordingly, the operation is easy and convenient, evading jeopardy of contamination human body or surrounding environment again.

[0037] Examples of the peroxide employed in the present invention include permanganate, sodium peroxide, various metal salts such as barium peroxide, zinc peroxide, cadmium peroxide, potassium peroxide, calcium peroxide. chromium peroxide, etc., persulfate, hydrocen peroxide and ozone.

15 [0038] Preferred peroxides used as a favorable oxidizing agent are permanganate and persulfate.

[0039] Typical examples of permanganate include zinc permanganate, cadmium permanganate, potassium permanganate, cadmium permanganate, silver permanganate, silver permanganate, silver permanganate, manganate, manganate,

hydrogen persulfate potassium, lead persulfate and rubidium persulfate, however, persulfates such as ammonium persulfate, sodium persulfate and potassium persulfate are particularly preferable as an oxidizing agent. They may be used alone or in combination of two or more kinds thereof. Enthermore, the use amount of the peroxide is preferably 100-fold mole or greater, more preferably selected within a range from 10<sup>4</sup>-fold mole to 10<sup>9</sup>-fold mole to 10<sup>9</sup>-fold mole and the most preferably selected within a range from 10<sup>4</sup>-fold mole on the basis of mole number of the hardy decomposable harmful substances adsorbed by solid in reference.

[0041] Regarding with the addition amount of peroxide, it is different depending on pH of wastewater to be treated, however, in the case where promoting only reaction, the addition may be conducted only taking an oxidizability of persulfuric acid into consideration.

[0042] Further, it is preferable for peroxide to be brought into contact with the solid in the state of dissolved in the wastewater in order for promoting the decomposition and furthermore, other oxidizing agent such as, for example, hydrogen peroxide or zone may be coexisted. Also, metal salt metallic oxide or so may be coexisted.

[0043] Still further, an organic solvent may be appropriately added into the reaction system in order to more effectively conduct the decomposition reaction. As the organic solvent, and vorganic solvent and sector selected from the group consisting of ketones having 3 to 6 carbon atoms, alcohols having 1 to 4 carbon atoms and carboxylate esters having 2 to 8 carbon atoms may be preferably employed. Among these, examples of the ketones having 3 to 6 carbon atoms include acetone, methyletis bettone, etherly tectone, etherly tectone, etherly tectone, etherly tectone, atoms include acetone, atoms include methylaclohol, ethyl alcohol, lspropanol, various butyl alcohols, ethylene glycol, propylene glycol, butylene glycol, etc. Furthermore, examples of the acohol atoms include methyl formate, ethyl formate, methyl acetate, ethyl acetate, methyl propionate, ethyl propanate, methyl butyrate, butyric ester, methyl acrylate, etc. Because these organic solvents have great affinity with water, they are capable of eluting the hardly decorposable harmly substances from inside of the solid to surface laver area of the solid.

[0044] Although persultates heatedly decompose and generate hydrogensulfate ion radical, sulfate ion radical and hydroxyl radical, and the radical further decomposes decomposable harmful substances such as dioxin, etc., it is preferable to agiltate after making the solid of the hardly decomposable harmful substances coagulated into a form of slurry in order to raise decomposition efficiency because the radical ejects electron by in a short time. The more violent the agitation is, the higher the probability that the radical and the hardly decomposable harmful substances come into contact with each other raises, and it is advantageous, however, it is preferable that the agitation is conducted within the range which does not become remarkably unfavorable in economic aspect regarding with a capacity of a decomposing container or viscosity of the slurry because the agitation has a limit.

[0045] Regarding with the reaction temperature for oxidative docomposition of the hardly decomposable harmful substances congulated to the solid with the use of peroxide, it is preferably from room temperature to 100 °C. The reaction temperature is more preferably within the range of from 40 °C to 100 °C. When the reaction temperature is lower than 40 °C, long time may be required for decomposition.

[0046] The higher the temperature for oxidative decomposition treatment, the faster the decomposition rate rses, however, the decomposition treatment at the temperature higher than the boiling temperature of water (when the salt concentration becomes high, the boiling temperature becomes higher than 100 °C) will require a pressure vessel, and accordingly, the decomposition treatment under atmospheric pressure at lower than the boiling temperature is preferble. Further, in the case where the decomposition treatment is conducted under the atmospheric pressure at 10 or 10

- temperature of boiling temperature or higher, it becomes necessary to provide waste gas treatment facility for the aspect of preventing secondary contamination because, at an elevated temperature, the hardly decomposable harmful substances such as dioxin evaporate together with evaporation of moisture.
- [0047] In an occasion of oxidative decomposition with the use of the above preferable persultate in the present invention, for the purpose of maintaining plt of 6 or higher, preferably 7 or higher by neutralizing the generating sulfuric acid, and for the purpose of adjusting the decomposition reaction, it is profitable to oxidatively decompose under the existence of hasis substance.
- [0048] With regard to the above basic substance, it is important to select a compound without being oxidized by persulfates and without inducing secondary contamination in an occasion of transferring the treated slurry towards an utimate disposal place such as a landfill, or of disposal. One or more kinds appropriately selected from hydroxide, oxides and slightly acid salt of alkali metal or alkaline earth metal; ammonia and organic base compounds is used as the basic substance.
- [0049] Preferable examples of the hydroxide of the alkall metal or the alkaline earth metals include hydroxides of sodium, potassium, calcium, etc., and preferable examples of the oxide of the alkall metals or alkaline earth metals include sodium oxide, potassium oxide, calcium oxide, barium oxide, sodium hyperoxide, potassium hyperoxide, calcium hyperoxide, barium hyperoxide, calcium hyperoxide, barium size oxide, barium oxide, potassium hyperoxide, calcium hyperoxide, barium size oxide, potassium oxide, potassium oxide, potassium oxide, potassium hydrogen carbonate, potassium hydrogen carbonate, potassium oxide, potassium oxi
- 20 [0050] These basic substances may be preparedly added into the solids, or may be added during the reaction little by little. In this way, melatishing pH of decomposing reaction solution to 6 or higher, preferably 7 or higher enables to employ a cheap reaction container made of iron without requiring to employ an expensive anticorrosion reaction container.
- [0051] In the present invention, healing method is not particularly restricted in an occasion of aciding heat, and any so one of an electric heater type, a heated water feed type, a steam suction type, a bolier type and so on may be employable, however, it is necessary to be careful about the heated water type that the water content should not become too much. Too much moisture content will reduce persulfate concentration for the reaction. With regard to treating time for oxidative decomposition, it is unable to settle in a wholesale manner because it depends on the treating temperature or other condition, however, it is usually from 10 minutes to about 50 hours.
- 30 [0052] Additionally, in the case where the hardly decomposable harmful substances strongly coagulate with the solid such as inorganic substances, it is desirable to oxidatively decompose the hardly decomposable harmful substances by preparedly bringing persulfates into contact with the solid and permeating them among the solid followed by heating or so.
- [0053] By the employment of the above oxidative decomposition treatment condition, favorable effects such as easy so operation, or evading jeopardy of contaminating human body or surrounding environment again based on the following merits: (f) it enables to reduce a use amount of persulfate; (2) it enables to shorten a reaction time; (3) it enables to lower a reaction temperature; (4) it enables to prevent corrosion of a reaction container; (5) it enables to carry out oxidative decomposition treatment with a cheap reaction container made of iron; and (6) it enables to oxidatively decompose the hardly decomposable harmful substances in a state of a solid.
- do [0054] Additionally, a slight amount of dioxin may be contained among a solution layer separated from a solid. In such a case, ozonization, ultraviolet irradiation treatment, catalyst treatment or activated carbon treatment may be applicable to the solution layer resultantly decomposing or adsorbing the remaining slight amount of dioxin.
- [0055] Next, the second aspect of the present invention provides a method of treating hardly decomposable harmful substance comprising (a) a step for adding adsorbent into wastewater containing a hardly decomposable harmful substance; (3) a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid; and (C) a step for decomposing the hardly decomposable harmful substance to with the separated solid.
  - [0056] Namely, it provides (a) a step for adding adsorbent into wastewater containing a hardly decomposable harmful substance instead of a step for adding coagulant into wastewater containing a hardly decomposable harmful substance in the first aspect of the present invention.
  - [0057] Additionally, (8) a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid, and (C) a step for decomposable harmful substance by bringing peroxide into contact with the separated solid may be carried out in the same manner as the forecomin (risk aspect of the orsent invention.)
- 55 [0058] Similarly as the step (A) in the present invention, it is desirable to preparedly adjust pH of the wastewater containing the hardly decomposable harmful substances within the range of from 6 to 12 before adsorbing the hardly decomposable harmful substances by adding the adsorbent to the wastewater as the step (a). When pH is lower than 6, pipes of the processing apparatus may corrode and when pH exceeds 12, neutralization of the wastewater at the

last step will become troublesome.

[0059] Carrying out the step (a), that is, adding the adsorbent into the pH adjusted wastewater after completing the above pH adjustment will adsorb the hardly decomposable harmful substances.

[0060] Regarding with a kind of adsorbent employed in the step (a), the same inorganic porous materials and organic porous materials which are the coagulation promoter employable in the foregoing first aspect may be employed appropriately. Further, Itlania (titan oxide) may be employed as the adsorbent. Concrete examples of the inorganic porous materials include activated calor, in exchange resin, etc. They may be used alone or in combination of two or more kinds thereof. Titania, activated caton, indicom earth and ion exchange resin are preferable among the adsorbents. Particularly, the activated carbon whose diameter of mesopore occupying the entire pore volume is 2 to 50 mm, macropore is 50 mm or greater, and whose pore volume is 0.15 fiterig or more, preferably 2.20 literig or more, and further preferably

0.25 ller/g or more is favorably applicable. [0061] Examples of the activated carbon not which the hardly decomposable harmful substances adsorbed include the following: (1) a mixture consisting of suspended matters and activated carbons having adsorbed dioxin obtained by cleaning gas exhausted from inclinerator plant facilities with scrubber and by conducting adsorptive treatment to the wastewater after the cleaning with the use of activated carbon; (2) activated carbons having adsorbed cloxin obtained by cleaning exhaust gas which generates when reproducing aluminum, zinc, iron, etc., with scrubber and by conducting adsorptive treatment to sutage after separating sold from the cleaning water and to wastewater after removing sludge with the use of activated carbon; (3) a mixture consisting of suspended matters and activated carbons having adsorbed substance obtained by conducting adsorptive treatment to wastewater from such as chloroethylinen production plant or apoxy resin production plant with activated carbon; and (4) activated carbons having adsorbed substance obtained by conducting adsorptive treatment to phenols in alkali cleaning wastewater drained from production plant of various alkybenous with activated carbon.

[0082] The adsorbent such as diatom earth or ion exchange resin to which the hardly decomposable harmful substances were adsorbed are capable of reducing the content of hardly decomposable harmful substances of Ital eff to a range sufficiently safe for reuse by means of the above-mentioned decomposition technique. Therefore, after once using the adsorbents for adsorption of the hardly decomposable harmful substances, the adsorbents are capable of being repeatedly used without disposing promptly and until degraciation of their properties as adsorbent occurs. Accordingly, they enable to treat wastewater as closed system, they exhibit extremely high safety and they are economic. Further, even in the case where the adsorbents for the hardly decomposable harmful substances are finally disposed, It is possible to dispose after reducing residue amount of the hardly decomposable harmful substances sufficiently, without giving adverse effect to natural environment.

[0063] Furthermore, the present invention is applicable to solid being adsorbed by the hardly decomposable substances. Namely, by applying the step (C) in the foregoing first aspect to the solid onto which the hardly decomposable substances were adsorbed, a method for decomposing the hardly decomposable substances were so are be rowing.

[0064] The present invention is favorably applicable to solid being adsorbed by the hardly decomposable substances, for example, soll, sludge, adsorbent and incinerated ash being adsorbed by the hardly decomposable substances.

[0065] Examples of the soil being adsorbed by the hardly decomposable substances include contaminated soil caused by ground-water containing hardly decomposable substances or by leachaler form at Indfill. Examples of the sludge being adsorbed by the hardly decomposable substances include suspended matters being adsorbed by dioxin obtained by separating and collecting suspended matters in effluent drained from each facility in factory and piace of business, and sludge being adsorbed by dioxin obtained by separating and collecting sludge from cleaning liquid after cleaning, with the use of a scrubber, exhaust gas which generates in an occasion of renewing waste catalyst discharged from a cleaning apparatus for exhaust gas which generates in an occasion of renewing waste catalyst discharged from a cleaning apparatus for exhaust gas discharged from male Irefining factory, etc.

5 [0065] Examples of the adsorbent being adsorbed by the hardly decomposable substances include inorganic porous material, organic porous material, ittania (titania (titania), etc. Examples of the inorganic porous material include zeolite, diatom earth, activated day, etc.

[0067] Further, examples of the organic porous material include activated carbon, ion exchange resin, etc.

[0068] Still further, examples of the incinerated ash being adsorbed by the hardly decomposable substances include incinerated ash being adsorbed by the hardly decomposable substances discharged after incinerating general garbage or industrial, garbage such as normal family life base wastes or business base wastes.

[0069] The hardly decomposable substances adsorbed onto the above solid are able to be exidatively decomposed by bringing them into contact with peroxides of 100-10d mole or more to the hardly decomposable substances without carrying out desorption operation. However, before or after bringing them into contact with the above peroxide, a contact treatment of bringing them into contact with microorganism and/or enzyme, or a contact treatment of bringing them into contact with present processing them into contact with present prese

[0070] The treatment by the microorganism or the enzyme will be explained below.

[0071] With regard to the microorganism used in the pre-treatment, mold is preferably employed. Typical example

of the mold belongs to Trameles genus. Schizophylium genus, Phanerochaete genus. Bjerkandere genus, Irpex genus, Pleurolus genus, Myceliophthora genus, Lentinera genus, Pycnoporus genus, Lentinus genus, Rhizoctonia genus, Funalia genus, Morellus genus, Myceliophthora genus, Coprinus genus, Agaricus genus, Ploriota genus, Flammulona genus. Ganoderma genus, Dadelaicopsis genus, Favolus genus, Lyophyllum genus, Auroluaria genus, Giloeophyllum genus, Tyrophyllum genus, Myceliophthora genus, Flammulona genus, Formatinum genus, Myceliophthora genus, Reurospora genus, Scherolum genus, Heterobasidion genus, Agaricus, Scherolum genus, Cladorthinum genus, Graphyllum genus, Schizophyllum genus, Schizophyllum genus, Schizophyllum genus, Sphaeropasis genus, Eusarium genus, Trichoders genus, Botryis genus, Aspargillus genus, etc. Among these molds, woods decay fungus bolonging to Trametes genus, Schizophyllum genus, Phaenecchaets genus, Bjerkandera genus, Irpex genus, Pleurotus genus, Myceliophthora genus, Lentinera genus and Pycnoporus) genus are particularly preferable.

[0721] On the other hand, at least one kind particularly selected from among cellobiose dehydrogenase, xanthine oxidase, laccase, lignin peroxidase and manganese peroxidase is preferably employed as the enzyme. With repart to the enzyme, those enzyme either which mitorograpism ejected after production or which are contained in natural product each separated from incubation broth and natural product with the use of ion exchange resin or so may be employed, and a mixture of visible fungus of microorganism and enzyme may be employed, in the case where the enzymes are employed, it is effective that, in an occasion of contacting with hardly decomposable substances, the enrolled be conducted either under the existence of electron donor or hydrogen peroxide or under the existence of microorganism or enzyme each producing hydrogen peroxide or compounds such as 1-hydroxyberotrazole and so on or aniline-based compounds such as 2,2- azino bis (3-ethylbenzthiazoline-6-sulfonic acid) are preferable to add a mediator in order to reveal that satisfy ultimately. Pehnolic compounds such as 1-hydroxyberotrazole and so on or aniline-based compounds such as 2,2- azino bis (3-ethylbenzthiazoline-6-sulfonic acid) are preferable to add was the mediator.

[0073] With regard to a method for incubating the mold, the same method for usually incubating microorganism may be applicable. For example, in a little incubation, incubating at the temperature of around 10 to 50 °C for 5 to 30 days on malt-yeast incubation medium may be appropriate. Further, in the case of incubating in large quantities, either a liquid incubation with the use of a tank or a solid incubation with the use of solid component of plant derivation such as sawdust, all graining of wheat or barley, bran or so, or with the use of inorganic porous carrier impregnated with sugar, nitrogen, phosphorus, mineral and so on may be conducted under the above condition. In this occasion, when the incubation temperature is lower than 10 °C , propagation of microorganism becomes slow and the amount of emitting enzyme decreases. Further, when the incubation temperature exceeds 50 °C, there will be a fear that growing proliferation of microorganism becoming slow. Moreover, it is preferable to adjust pH during incubation to 3 or higher. and further; it is more preferable to adjust pH to 3.5 or higher. When pH during incubation is lower than 3, the amount of emitting enzyme from microorganism becomes slow and it falls outside the range of optimum pH of enzyme. In this condition, corrosion will appear at inside of incubation container made of iron. Furthermore, in the incubation of the microorganism, a bacterial concentration of incubated microorganism might fall 1 × 10<sup>2</sup> cfu (a colony formation unit) per plant organic matter dry weight 1g or greater, preferably within a range of from  $1 \times 10^2$  to  $1 \times 10^9$  cfu, and more preferably within a range of from 1 × 103 to 1 × 107 cfu. Moreover, on the occasion of incubation of the mold, either mycellum or spore may be employed, however, mycellum is usually employed because of easiness in incubation.

[0074] When the microorganism and/or the enzyme is brought into contact with the hardly decomposable substances adsorbed to solid, it is preferable to bring the microorganism and/or the enzyme into contact under an aseptic condition after a sterilization treatment of saprophyte adhering to the solid beforehand. Various saprophytes usually habitate among the solid to which the hardly decomposable substances are adsorbed, and accordingly, by conducting sterilization treatment to the saprophytes, a favorable condition for a growth of useful microorganism after inoculating useful microorganism will be arranged.

[0075] With regard to a method for the sterilization treatment, a heat treatment method, a chemical treatment method or a physical treatment method may be applied. In the case where the heat treatment method is employed for conducting the sterilization treatment to the saprophytes, the treatment might be conducted at the temperature of around 80 to 125 °C. Although a heat treatment time is different depending on the treatment temperature, it might fall within a range of from 2 seconds to around 6 hours. Because almost all the saprophytes extinct at the temperature of 125 °C, the heat treatment does not to be conducted at the temperature exceeding it.

50 [0076] Further, in the case where the chemical treatment method is employed for the sterilization treatment, ctryl alcohol, diethyl discribonate, hydrogen peroxide, persulfate, hypochlorous acid, hydrochioric acid, ethylene oxide, ozone or chloropicirn is employable as a sterilization treatment agent. The treatment agents may be directly used; however, a solution obtained by using diluents such as water may be used. For example, in the case where ethyl alcohol is employed, an aqueous solution of it with a concentration of around 60 to 100 g/100milliliter is preferably sued. Further, in the case where hydrogen peroxide is employed, an aqueous solution of it with a concentration of less than 30 g/100milliliter perferably used, and a mixed aqueous solution with ethyl alcohol may be used. With regard to the physical treatment method, a method in accordance with an ultraviolet irradiation treatment is preferably used. [0077] Moreover, in an occasion of the decomposition treatment of bringing hardly decomposable substances ad-

sorbed to solid into contact with microorganism and/or enzyme, a propagation of saprophytes may be suppressed by urging a useful microorganism to grow selectively, aside from the stentization treatment. For example, in the case where wood decay fungus are employed as the microorganism, a use of a cellulose source as a carbon source; perfererably a use of carboxymethyl cellulose, water-soluble cellulose ether, phosphorylation cellulose and so on which are each soluble cellulose source as source respectively is capable of suppressing propagation of saprophytes because these carbon sources do not contribute to propagate saprophytes.

[0078] In the case where the hardly decomposable substances adsorbed to the solid sterilized by the heat treatment method, chemical treatment method or physical treatment method are decomposed being brought into contact with microorganism, a method of incubating the microorganism under the existence of the solid is desirable.

[0079]. With regard to a condition in the occasion of conducting decomposition reaction of hardy decomposable substances by the microorganism, it is similar with the growth condition of the microorganism. Themsely, the temperature is about 10 to 50 °C, preferably 15 to 35 °C, and pH is 3 or higher, preferably 3.5 or higher. Further, because the above mod is aerobic microorganism, the decomposition reaction should be conducted while supplying a little amount of gas containing enzyme, preferably air to a decomposition reaction device. As mentioned above, the incubation of microorganism under the existence of the hardy decomposable substances adsorbed to the solid will contribute to advance the decomposition reaction of the hardy decomposable substances because the incubation emits an enzyme capable of decomposing hardy decomposable substances and a radical capable of decomposing hardy decomposable substances and a radical capable of decomposing hardy decomposable substances with the decomposition that the decomposable substances with the decomposable substances and a radical capable of decomposing hardy decomposable substances with the decomposable with the decomposable with the decomposable with the decomposable with the decompos

[0080] Furthermore, in order to conduct the decomposition reaction more effectively, it is preferable to carry out the decomposition reaction under the existence of untifition source for the mold employed in the reaction. There are various substances as the nutrition source, and examples include saccharides such as glucose or so, carbon source such as potato extract or molasses, nitrogen source such as ammonium salt or urea, and water-soluble nutrition source such as corn steep liquor, meet extract, yeast extract or peptione. Moreover, corns such as barley and wheat, rice or corn; and their side-products such as bran, rice bran, cone broth. Okara and so on may be used as the nutrition source is Still further, wood chips. (they of coconut, skin of citrus or porous citry mineral may be added. With regard to the advanced to a mount of the nutrition source, in an occasion that the nutrition source is water-soluble and easy to be adsorted the solid. In a eddition amount of the nutrition source is solid.

and hardly adsorbed to the adsorbent, the addition amount is about 0.001 to 10 % by weight to the adsorbent. [0081] Still further, an organic solvent may be appropriately added into the reaction system in order to more effectively conduct the decomposition reaction. As the organic solvent, any organic solvent selected from the group consisting of ketones having 3 to 6 carbon atoms, alcohols having 1 to 4 carbon atoms and carboxylate esters having 2 to 6 carbon atoms may be preferably employed. Typical examples of the ketones having 3 to 6 carbon atoms include acetone, butanone, diethyl ketone, methyl isobutyl ketone, etc. Typical examples of the alcohols having 1 to 4 carbon atoms include methylalchol, ethylalcohol, isopropyl alcohol, various butylalcohols, ethylene glycol, propylene glycol, butylene glycol, etc. Typical examples of the carboxylate esters having 2 to 6 carbon atoms include methyl formate, ethyl formate, methyl acetate, ethyl acetate, methyl propionate, ethyl propanoate, methyl butyrate, butyric ester, methyl acrylate, ethyl acrylate, etc. The above-mentioned organic solvents have great affinity with water, and accordingly, the hardly decomposable substances in the state of eluted from inside of the solid to surface layer of the solid are effectively brought into contact with the microorganism or the enzyme without decreasing activity of the microorganism and the enzyme. Furthermore, together with the organic solvent, an addition of solvent such as liquid n-paraffins, cycloparaffins, higher fatty acid esters at an ordinary temperature enables to prevent re-adsorption of the hardly decomposable substances to the inside of the solid. Typical examples of the solvent for preventing re-adsorption include n-decane, n-undecane, n-dodecane, n-tridecane, n-tetradecane, n-pentadecane, n-hexadecane, cyclooctane, cyclodecane, methyl oleate. ethyl oleate, methyl linoleate, ethyl linoleate, methyl linolenate, ethyl linolenate, etc.

5 [0082] In the decomposition method of hardly decomposable substances of the present invention, after the pretreatment of optionally bringing microorganism or enzyme into contact with the hardly decomposable substances adsorbed in solid, the hardly decomposable substances are oxidatively decomposable by contacting with peroxides. Further, it is possible to decompose the hardly decomposable substances by bringing them into contact with the microorganism or the enzyme after bringing them into contact with the peroxide process.

2 [0083] The contact between the hardly decomposable substances adsorbed to the solid and peroxides is executed in the state that the hardly decomposable substances were adsorbed to the solid without substantially passing through a desorption treatment to make them olute from the solid. In order to effectively conduct the decomposition of the hardly decomposable substances in the above, the addition amount of the peroxide might be settled to 100-fold mole or greater, preferably settled to from 10° to 10°-fold mole and more preferably settled to from 10° to 10°-fold mole on the basis of mole number of the hardly decomposable harmful substances adsorbed to the solid.

[0084] In the oxidative decomposition, at least one kind of peroxide selected from the group consisting of salt of peroxo acid, hyperoxide and organic peroxide is preferably employed.

[0085] The salt of peroxo acid in the present invention is not particularly specified as far as it is safely handled, and

examples include various compounds such as persulfate, permanganate, perborate, organic persait, etc. Specific compounds of the salt of peroxo acid include persulfates such as ammonium persulfate, sodium persulfate, potassium persulfate, hydrogen persulfate potassium, etc.; permanganates such as sodium permanganate, potassium permanganate, calcium permanganate, etc.; perborates such as sodium perborate, potassium perborate, etc.; organic persaits such as persectic acid sodium, porthoracio acid sodium, porthhalia caid sodium, etc.

[0086] The hyperoxides are not particularly restricted and examples include various compounds such as potassium hyperoxide, sodium hyperoxide, barium hyperoxide, magnesium hyperoxide, etc.

[0087] Further, the organic peroxides are not particularly restricted and examples are various compounds including tection peroxides such as acceptionate organic particular peroxide, acceptionation peroxide, etc.; diacyj peroxides such as acceptionationation peroxide, bearvayl peroxide, laurcyl peroxide, 3.5.5-trimethyl hexanov) peroxide, etc.; hydro peroxides such as bebulylhydro peroxide, bearvayl peroxide, 2.5-dimethylbexano-2, 5-dihydro peroxide, etc.; dialkyl peroxide such as di-bubyl peroxide, didurnly peroxide, 2.5-dimethyl-25-dil(-buby) peroxy) hexan, etc.; peroxide such as 1-1-bis-t-bubyl peroxyl-peroxide, didurnly peroxide, 2.5-dimethyl-25-dil(-buby) peroxy) hexan, etc.; peroxyl ketalas such as 1-1-bis-t-bubyl peroxyl-peroxide, peroxyl-pe

[0088] In the present invention, the above peroxide may be used alone or in combination of two or more kinds thereof. [0089] In the present invention, the hardly decomposable substances presented with hardly decomposable substances, microorganism or enzyme adsorbed to the solid are oxidatively decomposed by the above peroxides just in the state of being adsorbed to the solid without substantially eluting them. In the above decomposition, the hyperoxides are usable in the form of power, aqueous solution or organic solvent solution depending on their form such as soil, sludge or incinerated ash, or depending on the character of the hyperoxide. In the case where the hyperoxides are used in the form of the aqueous solution or in the form of the organic solvent solution, the concentration is preferred by 0.1 % by weight, a large quantity of solution containing the hyperoxide will be needed and resultantly cause an inconvenience that a treatment container onlarges.

[0090] With regard to the contact treatment between the hardly decomposable substances and the hyperoxides, an oxidative decomposition bringing them into contact at Pd 3 or higher, preferably at pl 4.3 for higher is advantageous. [0091] Further, the temperature of the oxidative decomposition treatment bringing powdery peroxides or solution peroxide into contact with the hardly decomposable substances either adsorbed to the solid, or pretreated with micrograpiane or enzyme might be appropriately settled depending on the kind of the peroxide user, however, it using falls within the range of 20 to 20° °C and preferably 80 to 180° °C. In the present invention, heating method is not particularly restricted in an occasion of adding heat, and any one of an electric heater type, a heated water feel perox, a steam suction type, a boiler type and so on may be employable, however, it is necessary to be careful about the heated water type that the water content should not become too much. Too much moisture content will reduce peroxide concentration for the neaction. With regard to treating time for oxidative decomposition, it is unable to settle in a whole-sale manner because it depends on the treating temperature, the kind of the peroxide used or so, however, it is usually from 10 minutes to around 30 days.

[0092] Additionally, in the case where the hardly decomposable substances strongly adsorb to the solid such as active carbons or so, it is desirable to oxidatively decompose the hardly decomposable substances by preparedly bringing peroxides into contact with the solid sufficiently and permeating them among the solid followed by heating or so. Organic peroxide which easily permeates into activated carbon is more preferable to the decomposition of hardly decomposable substances adorbed especially to the activated carbon.

[0093] Furthermore, in the decomposition of the hardly decomposable substances absorbed onto the activated carbons, heating after preparedly adsorbing hyperoxides such as potassium hyperoxide, sodium hyperoxide, barthyperoxide or so each dissolved into an organic solvent respectively onto the activated carbon is also efficience.

[0094] Additionally, when the solid onto which the hardly decomposable substances were adsorbed have a strong sticking property, for example, when sludge containing both incinerated ash discharged from incineration furnace and activated carbon are decomposed, powdering the solid with grinder and simultaneously adding and mixing powdery peroxides into the sludge followed by the heat treatment is advantageous.

50 [0095] The present invention shall be explained below in further details with reference to examples, but the present invention shall by no means be restricted by the following examples.

## **EXAMPLES**

55 Example 1 and Comparative Example 1

[0096] An aqueous solution of poly aluminum chloride as a coagulant was continuously added by injection in an amount of 1 ppm as poly aluminum chloride content into the wastewater containing dioxin flowing at a flow rate of 1m<sup>3</sup>/

hr, and the resultant solution was filtered for 1 hour through a solid-liquid separator having a capacity of 10 L and equipped with a bag filter having membrane area of 0.25 m<sup>2</sup>. After the filtration, a part of the coagula which stuck to filter was gathered by means of a spatula, mixed enough, and after drying, an amount of dioxin was measured and determined as a blank (Comparative Example 1).

[0097] Further, 1 kg of sodium persulfate, 600 g of sodium hydroxide were thrown into water in an amount of up to 10 L, which was reacted for 6 hours in the solid-liquid separator with the state that the coagula stock to bag filter, keeping the temperature within the range of from 80 to 95 °C. After the reaction, a part of the coagula which stuct to the filter was gathered by means of a spatula, mixed enough, and after drying, an amount of dioxin residue was measured (Example 1). The measurement results are shown in Table 1. The results verify that high decomposition ratio was exhibited about the coagular.

#### Table 1

	Sample Condition	Measured amounts of Dioxin (ng/gdry)	Decomposition ratio (%)
F	Comparative Example 1	5 0	-
Г	Example 1	5	90

[0098] The solid-liquid separator equipped with the bag filter was manipulated as follows. Explanation will be described referring to Ffig. 1. Fig. 1 illustrates a block diagram of a solid-liquid separator with a bag filter employed in Examples and Comparative Examples.

(1) Solid-liquid separation step (filtration concentration step)

[0099] Valves 1, 5 and 6 were opened and valves 2, 4, 7, 8 and 9 were closed. Valve 6 was filled with the solution and then, turned to a closed state. With the use of pump 12, the coagulant was injected into drain line 16 through valve 7 before entering into bag filter 10. This system was designed for introducing the wastewater into the solid-liquid separator utilitizing a fall of drain line 16.

(2) Step for decomposing hardly decomposable substance (reaction step)

[0100] Also with the use of pump 12, after opening valves 6 and 7 and closing other valves, a reagent was injected from a reagent tank by operating pump 12. After the injection of the reagent, closing valves 6 and 7 and opening valves 2, 3 and 4, further operating pump 13, the solution was cycled and heated by means of heat exchanger 14, and resultantiv reacted.

[0101] Additionally, reference numeral 11 indicates a pressure gauge and reference numeral 15 indicates a temperature controlling trap.

Example 2 and Comparative Example 2

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O1022 A suspension containing 250 g/L of diatore earth was poured in an amount of 1 L through valve 6 into solid-liquid soparator having a capacity of 10 L and equipped with a bag filter having a membrane area of 0.25 m², and as a result, the surface of the bag filter was preceated with diatore earth only by filtering the suspension. Subsequently, an aqueous solution of poly aluminum chloride was continuously added by injection in an amount of 1 ppm as poly aluminum chloride content into the wastewater containing dioxin flowing at a flow rate of 1 m³/nr for 3 hours, thereby filtered the resultant wastewater. After the filtration, a container of the bag filter was opened up and a part of a cake was sampled from the filter, and after drying, an amount of dioxin was measured and determined as a blank (Comparative Example 2).

[0103] Further, an aqueous solution combining 1.5 Kg/SL of sodium persulfate and 1 kg/SL of sodium hydroxide, an aqueous solution of sodium persulfate in an amount of 5 L were thrown in this order into the solid-fiquid separator with the state that the coagula stuck to bag filter. Then, keeping the temperature within a range of 70 to 80 °C, a decomposition reaction of the resultant solution was carried out for 7 hours. After the termination of the reaction, a part of the reactant was sampled and after drying, an amount of doxin residue was measured (Example 2).

5 Example 3 and Comparative Example 3

[0104] Comparative Example 3 was conducted in a similar manner as Comparative Example 2 except that an aque-

ous solution of poly aluminum chloride as a coagulant was continuously added by injection in an amount of 1 ppm as poly aluminum chloride content into the wastewater and an aqueous solution of polyferric sulfate was also continuously added by injection in an amount of 10 ppm as polyferric sulfate content into the wastewater (Comparative Example 3). [0105] Further, the decomposition reaction was carried out in the same mnner as Example 2 about the sample obtained in Comparative Example 3). The measured data about the amount of dioxin each in Comparative Examples 2 and 3 are shown in Table 2 respectively.

Table 2

Sample Condition	Measured amounts of Dioxin (ng/gdry)	Decomposition ratio (%)
Comparative Example 2	58. 6	
Example 2	1. 8	9 6. 9
Comparative Example 3	6 0. 7	-
Example 3	1. 2	98.0

Example 4 and Comparative Example 4

[016] Pouring 100 ml of water containing dioxin into two centrifuge tubes made of polytetrafluoroethylene each having volume of 300 ml, and adding poly aluminum chloride and ferric sulfate each as coagulants in an amount that they each correspond to 2 ppm and 10 ppm respectively, and then, they were left standing for 16 hours. Subsequently, centrifugal separation was conducted with acceleration of 8000 G for 10 minutes. The content of one centrifuge tube was divided into supermatant and sediment as in Comparative Example 4, and a dioxin concentration was measured about each of them. Adding 100 ml of delonized water, 10 g of ammonium persuitate and 6 g of sodium hydroxide into the sediment in the other centrifuge tube, they were reacted keeping the temperature within a range of 85 to 95 for 7 hours. The reaction was named as Example 4 and a dioxin concentration was measured about the reactant. The results are shown in Table 5.

Toblo

		iau	)Ie 3	
30	Sample Condition	Category	Measured amounts of Dioxin (ng / gdry)	Decomposition ratio (%)
	Comparative Example 4	Supernatant	Under analytical sensitivity	-
35		Sediment	14	
	Example 4	Sediment after decomposition	0. 11	9 9. 2

40 Examples 5 to 8 and Comparative Examples 5 to 8

[0107] Examples 5 to 8 and Comparative Examples 5 to 8 are carried out in the same manners as Example 4 and Comparative Example 4 except that the coegulants are replaced to various ferrous coagulants shown in Table 4. The measured data about the amount of dioxin are shown in Table 4.

Table 4

	Addition amour coagulant (Main		Amounts of Dioxin (ng / gdry)				
			Supernatant	Coagula (ng)	Decomposed substance (ng)		
Comparative Example 5	Ferric sulfate	60	Under analytical sensitivity	20.5	_		
Example 5					1.0		

5.5

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Table 4 (continued)

	Addition amour coagulant (Main		Amoun	Amounts of Dioxin (ng / gdry)		
			Supernatant	Coagula (ng)	Decomposed substance (ng)	
Comparative Example 6	Polyferric sulfate	60	Under analytical sensitivity	10.5	-	
Example 6	1		-	_	0.6	
Comparative Example 7	Ferric chloride	60	Under analytical sensitivity	19.0	-	
Example 7	1		-	•	1.0	
Comparative Example 8	Polyferric chloride	60	Under analytical sensitivity	14.5	-	
Example 8			_	_	0.7	

### 20 Examples 9 to 12 and Comparative Examples 9 to 12

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[0108] Examples 9 to 12 and Comparative Examples 9 to 12 are carried out in the same manners as Example 4 and Comparative Example 4 except that the coagulants are combinedly employed of various kinds shown in Table 5. The results are shown in Table 5.

Table 5

				ble 5			
		Combination and amount of coagul		Amounts of Dioxin (ng / gdry)			
30				Supernatant	Coagula (ng)	Decomposed substance (ng)	
	Comparative Example 9	Ferric sulfate Polyacrylic acid	3 0 ppm 3 ppm	Under analytical sensitivity	17. 5	-	
15	Example 9	] ' '		_	_	0.06	
	Comparative Example 10	Polyferric sulfate	2 0 ppm	Under analytical sensitivity	15. 5	-	
10		Polyaluminum chloride	5 ppm				
-		Polyacrylic acid	1 ppm				
	Example 10			_	_	0.03	
5	Comparative Example 11	Ferric chloride Polyaluminum chloride	4 0 ppm 4 ppm	Under analytical sensitivity	13. 5		
	Example 11	Polyacrylic acid	3 ppm	_	-	0.7	
io	Comparative Example 12	Polyferric chloride	3 0 ppm	Under analytical sensitivity	9. 9	-	
	Example 12	Polyaluminum chloride	3 ppm	_	_	0.3	

<sup>55</sup> Examples 13 and 14, Comparative Examples 13 and 14

[0109] Fig. 1 illustrates a block diagram of a solid-liquid separator with a bag filter employed in Examples 13, 14, Comparative Examples 13 and 14.

[0110] A suspension containing 250 g/10L of diatom earth was poured through valve 6 into a solid-liquid separator equipped with a bag filter having a membrane area of 0.25 m<sup>2</sup>, and as a result, the surface of the bag filter was precoated with diatom earth only by filtering the suspension. Subsequently, the wastewater containing dioxin was flown at a flow rate of 1 m3/hr for 3 hours, thereby filtered the wastewater. After the filtration, a container of the bag filter was opened up and a part of a cake was sampled from the filter, and after drying, an amount of dioxin was measured and determined as a blank (Comparative Example 13).

[0111] Further, an aqueous solution combining 1.5 kg/5L of sodium persulfate and 1 kg/5L of sodium hydroxide, an aqueous solution of sodium persulfate and an aqueous solution of sodium hydroxide were thrown in this order into the solid-liquid separator. Then, keeping the temperature within a range of 70 to 80 °C, a decomposition reaction of the resultant solution was carried out for 7 hours. After the reaction terminated, a part of diatom earth was sampled, sufficiently agitating, and after drying, an amount of dioxin residue was measured (Example 13),

[0112] Subsequently, inside of the filter was washed by reverse flow and the decomposed substances were removed outside the system, repeating similar operations with the use of the same filter. After the filtration, a part of the diatom earth was sampled from the filter, sufficiently agitating, and after drying, an amount of dioxin was measured and determined as a blank (Comparative Example 14).

[0113] Further, an aqueous solution combining 1.5 kg/5L of sodium persulfate and 1 kg/5L of sodium hydroxide, an aqueous solution of sodium persulfate and an aqueous solution of sodium hydroxide were thrown in this order into the solid-liquid separator. Then, keeping the temperature within a range of 70 to 80 °C, a decomposition reaction of the resultant solution was carried out for 7 hours. After the reaction terminated, a part of diatom earth was sampled, sufficiently agitating, and after drying, an amount of dioxin residue was measured (Example 14).

[0114] The solid-liquid separation equipment equipped with the further bag filter was manipulated as follows. Explanation will be described referring to FIG. 1.

- (1) Solid-liquid separation step (filtration concentration step)
- [0115] Valves 1, 5 and 6 were opened and valves 2, 4, 7, 8 and 9 were closed. Valve 6 was filled with the solution and then, turned to a closed state. The adsorbent was injected through valve 6. This system was designed for introducing the wastewater into the solid-liquid separator utilizing a fall of drain line 16.
- (2) Step for decomposing hardly decomposable substance (reaction step)
  - [0116] Also with the use of pump 12, after opening valves 6 and 7 and closing other valves, a reagent was injected from a reagent tank by operating pump 12. After the injection of the reagent, closing valves 6 and 7 and opening valves 2. 3 and 4. further operating pump 13, the solution was cycled and heated by means of heat exchanger 14, and resultantly reacted.
  - [0117] Additionally, reference numeral 11 indicates a pressure gauge and reference numeral 15 indicates a temperature controlling trap.
- (3) Back wash inside the filter an
  - [0118] Valves 2, 3, 4, 5, 6 and 9 were opened and the other valves were closed. City water was connected to the line directly tied with valve 9, into which the water was injected. Valve 6 was closed after degassing.
- [0119] The measured data about the amount of dioxin each in Examples 13. 14, Comparative Examples 13 and 14 are shown in Table 6 respectively. From the above result, it was recognized that the reaction advances in closed system without manual operation and that the filter is reusable.

Γaŀ	

Sample Condition	Measured amounts of Dioxin(ng/g dry)	Decomposition ratio (%)
Comparative Example 13	61. 1	-
Example 13	2. 9	95. 2
Comparative Example 14	61. 0	1
Example 14	2. 5	95. 9

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Examples 15 to 20. Comparative Example 15

[0120] Suspended substances accumulated in cleaning water for exhaust gas of incineration plant were precipitated and then, concentrated by suction filtration with the used of a filter of 0.4 µm mesh, thereby gathered sudge. A dioxin concentration in this sludge in moisture content 62 % was measured and it was 45 no /a div sludge.

[0121] Throwing 20 g of the sludge into a container made of polytetralluorealtylene having a capacity of 100 ml and then after adding various kinds of salt of peroxo acid powder, stirred at the rate of 10,000 rounds per minute for 3 minutes at room temperature with the use of a high speed mixer. The container made of polytetralluorealtylene was placed into an autoclave, and it was heated for a predetermined time at various temperatures. Afterwards, dioxin concentrations about total amounts of samples in the polytetralluorealtylene container were measured.

[0122] Operating condition and results in each Examples and Comparative Examples are shown in Table 7. In Table 7, the addition amounts of peroxide equivalent to what numbers of fold mole per dioxin among studge were also shown. Further, the dioxin concentration was expressed with concentration corresponding to dry solid. The above descriptions are similarly applied to each following Examples and Comparative Examples. Additionally, various kinds of dioxins are contained in studge, however, calculation of the folded mole was carried out with the use of \$22 as a molecular weight of dioxin, which is a molecular weight of 2.3.7,8-letrachlorodibenzo oxine having the greatest toxicily. The above description is also similarly apoliced to each following Examples 21 to 40 and Comparative Examples 16 to 26.

Table 7

				Table	7		
20		Peroxides			Treated Temperature (°C)	Treated Time (hr)	Amount of Dioxin (nanogram / g sludge)
25		Kind	Addir	tion amount			
			(g)	(fold mole)			
	Example 15	Ammonium persulfate	10	4.2x10 <sup>7</sup>	100	5	0.8
30	Example 16	Ammonium persulfate	5	2.1x10 <sup>7</sup>	100	5	3.8
	Example 17	Ammonium persulfate	10	4.2x10 <sup>7</sup>	100	1	5.7
35	Example 18	Ammonium persulfate	10	4.2x10 <sup>7</sup>	80	15	16
	Comparative Example 15	-	-	-	100	5	43
40	Example 19	Sodium persulfate	10	4.2x10 <sup>7</sup>	100	5	6.3
	Example 20	Potassium persulfate	10	3.5x10 <sup>7</sup>	100	10	8.6
45		Amount o	f dioxin a	among the trea	ted sludge	•	45

Examples 21 to 23, Comparative Example 16

[0123] Reclaimed waste gas of complex catalyst was cleaned with a scrubber, and solid were separated from the cleaning water with the use of rotary flat membrane separator, followed by concentration with the use of a centrifugal separator.

[0124] Throwing 20 g of the solid (moisture content 67 %) into a container made of polyletrafluoroethylene having a capacity of 100 ml and then after adding persoxo acid at room temperature, stirred at the rate of 12,000 rounds per minute for 3 minutes with the use of a high speed mixer. The container made of polytetrafluoroethylene was placed 55 ml oan autociave, and it was heated for a predetermined time at various temperatures. Afterwards, dioxin concentrations about total amounts of samples in the polytetrafluoroethylene container were measured.

[0125] Operating condition and results in each Examples and Comparative Examples are shown in Table 8.

#### Table 8

5		Peroxides		Treated Temperature (°C)	Treated Time (hr)	Amount of Dioxin (nanogram / g sludge)	
		Kind	Addi	tion amount			
			(g)	(fold mole)			
10	Example 21	Ammonium persulfate	10	4.5x10 <sup>6</sup>	100	15	22
	Example 22	Sodium persulfate	10	4.4x10 <sup>6</sup>	100	15	28
15	Example 23	Potassium persulfate	10	3.8x10 <sup>6</sup>	100	15	36
	Comparative Example 24	-	-	-	100	15	470
20		Amount	of dioxin	among the tre	ated solid	•	470

Examples 24 to 27 and Comparative Example 17 to 21

[0126] Sludge accumulated at the bottom of deposition site of leachate from trash incinerated sah reclaimed land were gathered and concentrated at excleration of 8 000 G with he use of a continuous centrifugal separator. Through 20 g of the sludge (moisture content: 72 %) into a container made of polysterafluoroethylene having a capacity of 100 ml and then after adding various kinds of salt of peroxo acid, stirred at the rate of 10,000 counds per minute for 3 minutes at room temperature with the use of a high speed mixer. The container made of polysterafluoroethylene os as placed into an autoclave, and it was heated for a predetermined time at various temperatures. Afterwards, dioxin concentrations about total amounts of samples in the polyterafluoroethylene container were measured.

[0127] Operating condition and results in each Examples and Comparative examples are shown in Table 9.

#### Table 9

				lable a			
35			Peroxides			Treated Time (hr)	Amount of Dioxin (nanogram / g)
		Kind	Additio	n amount			
40			(ng)	(fold mole)			
	Example 24	Ammonium persulfate	2x10 <sup>9</sup>	1.3x10 <sup>7</sup>	100	15	15
45	Example 25	Ammonium persulfate	8x10 <sup>9</sup>	5.3x10 <sup>7</sup>	100	15	8.6
	Example 26	Ammonium persulfate	1.5x10 <sup>10</sup>	1.0x10 <sup>8</sup>	100	15	1.8
50	Example 27	Ammonium persulfate	2x10 <sup>10</sup>	1.3x10 <sup>8</sup>	90	15	16
	Comparative Example 17	-	-	-	100	15	37
55	Comparative Example 18	Ammonium persulfate	7.5x10 <sup>2</sup>	5.0	100	5	41
	Comparative Example 19	Ammonium persulfate	7.5x10 <sup>3</sup>	5.0x10	100	15	36

Table 9 (continued)

		Peroxides		Treated Temperature (°C)	Treated Time (hr)	Amount of Dioxin (nanogram / g)
	Kind	Additio	Addition amount			
		(ng)	(fold mole)			İ
Comparative Example 20	Ammonium persulfate	1.1x10 <sup>4</sup>	7.5x10	100	15	38
Comparative Example 21	Ammonium persulfate	1.4x10 <sup>4</sup>	9.0x10	100	15	41
	Amour	nt of dioxin an	nong the treate	ed sludge		38

Examples 28 to 30 and Comparative Example 22

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[0128] Reclaimed waste gas of complex catalyst was cleaned with a scrubber, and solid were separated from the cleaning water with the use of rotary flat membrane separator, followed by concentration with the use of a centrifugal separator.

[0129] Throwing 20 g of the solid (moisture content 67 %) into a container made of polyterafluoroethylene having a capacity of 100 ml and then after adding peroxo acid at room temperature, stirred at the rate of 12,000 rounds per minute for 3 minutes with the use of a high speed mixer. The container made of polyterafluoroethylene was placed into an autocalwe, and it was heated for a predetermined time at various temperatures. Afterwards, dioxin concentrations about total amounts of samples in the polyterafluoroethylene container were measured.

[0130] Operating condition and results in each Examples and comparative Examples are shown in Table 10.

Table 10

				IGDIC	10			
30		Peroxides			Treated Temperature (°C)	Treated Time (hr)	Amount of Dioxin (nanogram / g)	
		Kind	Addi	tion amount				
35			(g)	(fold mole)				
35	Example 28	40 % Sodium permanganate	6	7.2x10 <sup>5</sup>	70	15	85	
	Example 29	Potassium hyperoxide	5	3.0x10 <sup>6</sup>	100	15	104	
40	Example 30	70 % tertiarybutyl hydroperoxide	5	1.7x10 <sup>6</sup>	45	5	142	
15	Comparative Example 22	-	-	-	100	15	360	
		Amount of dioxin among the treated solid						

Examples 31 to 33 and Comparative Example 23

[0131] Soils were collected from the depth of 5 cm from the top of a surface layer of soil at a distance of 50 m from incineration plant for garbage. One hundred grams (water: 34 %) of the collected soils were placed into glass-made conical flask having a capacity of 500 ml.

[0132] On the other hand, 5 g of sodium permanganate with 40 % concentration was thrown into 50 ml of water, and the resultant solution was poured into the conical flask thereby forming slurny, followed by stirring at room temperature by means of an agitating rod for 30 minutes. Subsequently, the conical flask was heated at various temperatures while stirring. Afterwards, the total amount of the soils among the flask was analyzed.

[0133] Further, a section without adding sodium permanganate was provided as Comparative Example 23.

[0134] Operating condition and results in each Examples and Comparative Examples are shown in Table 11.

#### Table 11

	Heating temperature (°C)	Heating time (hr)	Amount of Dioxin (nanogram / g)			
Example 31	30	5	10			
Example 32	45	5	8.7			
Example 33	70	5	6.6			
Comparative Example 23	70	5	32			
Amount o	f dioxin among the treated so	oil .	35			

Examples 34 and 35. Comparative Examples 24 and 25

[0135] Dioxins among soil were decomposed with the use of sodium permanganate in a similar manner as Example 32 except that 1 kg of soil was thrown into a conical flask having a capacity of 5 L and that 10 gol 40 % sodium permanganate dissolved into 50 m of water and then, the resultant solution was added into the flask.

[0136] Afterwards, taking out the soil from the conical flask, transferred them in the amount of 50 g respectively into conical flasks each having capacities of 300 ml.

[0137] Adding 4 g of beech sawdust (moisture content: 16 %) and 10 ml of water into each flask, the contents were mixed.

The flasks were placed into an autoclave and they were sterilized at the temperature of 121 °C for 15 minutes.

[0138] On the other hand, beech sawdust in an amount of 7 part by weight, bran in an amount of 3 part by weight and CMC in an amount of 0.5 part by weight were mixed, and after adjusting the moisture content to 65%, the resultant mixture was extrusion molded into pellets having a size of 10 mm in diameter.

[0139] The pellets were sterilized in an autoclave at the temperature of 121 °C for 20 minutes and then, using them as incubation mediums, wood decay fungus were inoculate into them, followed by incubating at the temperature of 30 °C for 14 days.

[0140] The pellets were added into the foregoing dioxins-contaminated soil and the resultant soil was incubated at the temperature of 30°C for 30 days. Afterwards, an amount of dioxin in the flask was determined by content analysis. [0141] For the purpose of comparison, a section without adding wood decay fungus and a section adding excess sludge of activated sludge instead of wood decay fungus were provided.

[0142] Operating condition and results in each Examples and Comparative examples are shown in Table 12.

#### Table 12

	Microorganism	Amount of dioxin residue (nanogram / g)	
Example 34	Schizophillum commune IF06505	7.3	
Example 35	Gloeophyllum trabeum IF06268	6.3	
Comparative Example 24	Without inoculation	10	
Comparative Example 25	Excess sludge	11	

45 Examples 36 to 40. Comparative Example 26

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[0143] Reclaimed waste gas of complex catalyst was cleaned with a scrubber, and the resultant wastewater was passed through an advivated carbon cistema thereby adsorbing dioxins. Taking out the activated carbon from the cisterna, each 50 g (moisture content: 53 %) were thrown into polytetrafluoroethylene containers having capacities of 100 ml and 50 ml of water was further added respectively. The polytetrafluoroethylene containers were placed into iced water, and affer adding organic peroxide, peroxides were made to adsorb onto the activated carbon by stirring the contents of the containers for 24 hours.

[0144] Afterwards, placing the polyterafluoroethylene containers into incubators of various temperature, dioxins adsorbed onto the activated carbon were decomposed by reciprocate-shake of the incubators. Afterwards, a total amount of dioxin in the polyterafluoroethylene container was determined. Additionally, the calculation of the amount was carried out on the condition that the specific gravity of di-tertiarybutyl peroxide is 0.79, the bulk density and the concentration of dicumyl peroxide powder are 0.58 and 40 % respectively.

[0145] Operating condition and results in each Examples and Comparative Examples are shown in Table 13.

Table 13

				TCLOT				
5		Peroxides		Heated temperature (°C)	Decomposition time (hr)	Amount of Dioxin (nanogram / g )		
	Example 36	Ditertiarybutyl peroxide	250	5.1x10 <sup>10</sup>	30	5	6.7	
10	Example 37	Ditertiarybutyl peroxide	250	5.0x10 <sup>10</sup>	45	5	3.8	
	Example 38	Ditertiarybutyl peroxide	250	4.9x10 <sup>10</sup>	80	5	29	
15	Example 39	Dicumyl peroxide	250	5.0x10 <sup>10</sup>	30	5	8.2	
	Example 40	Dicumyl peroxide	250	5.1x10 <sup>10</sup>	45	5	3.4	
20	Comparative Example 26	-	-	-	80	5	34	
		Amount of dioxin among treated activated carbon						

Examples 41 to 45 and Comparative Examples 27 to 28

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[0146] Soil contaminated by heavy oil was vacuum concentrated at the temperature of 120 °C and removed light hydrocarbon from the soil. Fifty grams of the soil was dividedly taken into a glass container, and an aqueous solution of peroxa acid was further added into the container, then, after sufficiently stirring the content of the container, it was placed into an autoclave, which was treated at various temperatures for a predetermined time. Afterwards, polycyclic aromatic hydrocarbon in the soil was extracted and determined. Operating condition and results in each Examples and Comparative Stamples are shown in Table 14.

5		3/kg of soil)	۵	40	12	œ	14	os .	76	17	88	
		Hardly decomposable substance (mg/kg of soil)	O	61	31	26	31	23	128	127	130	
10			8	72	38	24	48	32	133	128	135	
15		Hardly deco	∢	63	21	F	24	19	115	107	115	
20			Time (hr)	ις.	so.	υ O	ω.	ဖ	ω.	G.		
25	14	Heating	Temperature (°C)	100	100	120	100	100	120	120		D : Benz (a) anthrathene
30	Table 14		Addition amount (ml)	1000	100	100	100	100	100	10,000	mong treated soil	
40		Peroxides	Concentration of solution (%)	0.2	2	2	4	2	0	0.02	Hardly decomposable substance among treated soil	uoranthene C:Chrysene
45			Kind	E a	mn je	m eg	E e	yanate		En es	Hardly dec	B : Benz (b) fluoranthene
50			×	Ammonium persulfate	Ammonium persulfate	Ammonium	Potassium persulfate	Sodium permanganate		Ammonium persulfate		
55				Example 41	Example 42	Example 43	Example 44	Example 45	Comparative Example 47	Comparative Example 48		A: Benz (a) pyrene

#### Examples 46 to 49

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[0147] Dissolving 5,000 µg of bisphenol A, 5,000 µg of bisphenol S, 2,000 µg of cotyl phenol, 2,000 µg of nonyiphol and 2,000 µg of pentachlorophenol into 50 ml of ethanol, the resultant solution was poured into a glass spray device. The ethanol solution was sprayed onto 1 kg (moisture content: 16 %) of reddish brown soil. After further pouring 50 ml of ethanol into the spray device, the resultant co-cleaning liquid was sprayed onto the soil. The soil was vacuum dried until moisture content reached 8 %, and the compound was made to adsort on to the soil.

[0148] Transferring 25 g of the soil into a container having a capacity of 250 ml and after adding 25 ml of aqueous solution of various kinds of peroxo acid, the resultant solution was recaled at various temperatures in an autoclave. [0149] Operating condition and results in each Examples are shown in Table 15.

5		soil)	Pentachlorophenol		220		360		870	490	2010							
10		Hardly decomposable substance (mg/kg of soil)	Hardly decomposable substance (mg/kg of	Nonylphenol		150		120		410	140	1780						
15				Hardly decomposable subst	Hardly decomposable subst	mposable subst	nposable subst	nposable subst	nposable substa	Octylphenol		330		520		280	160	1830
20						Bisphenol S		45		98		1300	800	4600				
25	Table 15		Bisphenol A		280		089		2300	092	4900							
30	Table	Treated Time (hr)			8		8		80	80	e treated soil							
40		Treated Temperature (°C)			100		100		46	100	Concentration of the hardly decomposable substances among the treated soil							
45		kides	Concentration	(J/B)	20		20		50	90	ecomposable sub							
50		Peroxides	Kind		Sodium	permanganate	Potassium	permanganate	Ammonium persulfate	Ammonium persulfate	on of the hardly d							
55					Example 46		Example 47		Example 48	Example 49	Concentrati							

#### Comparative Examples 29 and 30

[0150] Comparative Examples 29 and 30 were carried out in similar manners as Examples 48 and 49 except that the concentrations of the aqueous solution of ammonium persulfate were changed.

[0151] Operating condition and results in each Comparative Examples are shown in Table 16.

5		e e	Pentachloro phenol		1980	1910	2010		
10		Hardly decomposable substance (mg/kg of soil)	Nonylphenol		1730	1650	1780		
15			Hardly decomposable substr	Hardly decomposable subst	Octylphenol		1850	1690	1930
20					Bisphenol A Bisphenol S		4300	4200	4600
25	"		Bisphenol A		4800	4200	4600		
30	Table 16	Treated Time (hr)			80	∞	treated soil		
35		Treated Temperature (°C)			45	100	ances among the		
45		sep	Concentration	(1/6)	0.3	0.3	omposable substa		
50		Peroxides	Kind		Ammonlum persulfate	Ammonium persulfate	Concentration of the hardly decomposable substances among the treated soil		
55					Comparative Example 29	Comparative Example 30	Concentration		

#### Examples 50 to 52

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[0152] Reclaimed waste gas of complex catalyst was cleaned with a scrubber, and solids were concentrated in the cleaning water with the use of a continuous centrifugal separator.

[0153] Further, a solid cake was formed by applying the concentrates to the centrifugal separator.

[0154] Throwing 20 g of the solid cake (moisture content; 76 %) into a container made of polyfetrafluoroethylene having a capacity of 10 ml and then after adding 10 % sodium hyproxidos doubtion, sirred at the rate of 12,000 rounds per minute for 3 minutes with the use of a high speed mixer. Afterwards, adding 50 ml of 10 % aqueous solution of armmonium persulfate, the resultant solution was further stirred for 3 minutes with the use of the high-speed mixer. The container made of polytetrafluoroethylene was placed into an autoclave, and it was heated for a predetermined time at various temperatures. Afterwards, dioxin concentrations about total amounts of samples in the polytetrafluoroethylene container were measured.

[0155] Operating condition and results in each Example are shown in Table 17.

Table 17

	Addition amount of	Treated	Treated	Amount of	pH after
	sodium hydroxide solution	Temperature	Time	Dioxin	treatment
	(ml)	(°C)	(hr)	(nanogram / g )	
Example 50	8.75	100	15	4.7	4.5
Example 51	17.5	100	15	3.6	<12
Example 52	0	100	15	6.2	<1
Amou	nt of dioxin among tr	eated solid cake		60	

#### 35 Comparative Examples 31 to 33

[0156] Comparative Examples 31 to 33 were carried out in the same manners as Examples 50 to 52 except that city water was added instead of the aqueous solution of ammonium persulfate for the purpose of reference. Operating condition and results in each Examples and Comparative Examples are shown in Table 18.

Table 18

			Iable 18		
15		Addition amount of sodium hydroxide solution (ml)	Treated Temperature (°C)	Treated Time (hr)	Amount of Dioxin (nanogram / g)
15	Comparative Example 31	8.75	100	15	55
	Comparative Example 32	17.5	100	15	55
iO	Comparative Example 33	0	100	15	60
		Amount of dioxin amou	ng treated solid cake		60

[0157] As the above results, the decomposition advanced enough even under the condition of from slightly acidic to alkalescence. The fact that the reaction advances under alkali condition using peroxo acid is not known and to a daylantage in the selection, maintenance and management of materials because corrosion of the decomposition container may be reduced remarkably.

#### Example 53 and Comparative Example 34

9 [0158] Pouring 100 ml of water containing dioxin into 2 centrifuge tubes made of polyterafluoroethylene each having volume of 300 ml, and adding titian oxide (eavilable from Ishihara Sangyo Ksisha. Ltd., trade name. ST-01) as an adsorbent in an amount that each correspond to 400 pmm respectively, and then, they were agitated for 12 hours by means of a serve shaker. Subsequently, centrifugal separation was conducted with acceleration of 8000 G for 10 minutes. The content of 1 centrifuge tube was divided into supermatant and sediment as in Comparative Example 34, 40 and a dioxin concentration was measured about each of them. Adding 100 ml of deionized water, 1 go is sodium persulfate and 0.5 g of sodium hydroxide into the sediment in the other centrifuge tube, they were reacted keeping the temperature at 70 °C for 24 hours. The reaction was named as Example 53 and a dioxin concentration was measured about the reactant. The measurement results are shown in Table 19.

Table 19

Sample Condition	Category	Measured amounts of Dioxin (ng / pipe)	Decomposition ratio (%)	
Comparative Example 34	Supernatant	Under analytical sensitivity		
	Sediment	6. 8		
Example 53	Sediment after decomposition	0.7	90	

Example 54 and Comparative Example 34

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[0159] Pouring 100 ml of water containing dioxin into 2 centrifuge tubes made of polytetrafluoroethylene each having volume of 300 ml, and adding titan oxide (available from Ishihara Sangyo Kaisha, Ltd., trade name: ST-01) and an inorganic oxegulant (available from EARTH CLUSTER Incorporated, trade name: MOSSNITE) as adsorbents each in an amount that corresponds to 400 ppm and 100 ppm respectively, and then, they were agitated for 12 hours by means of a sieve shaker. Subsequently, centrifugal separation was conducted with acceleration of 8000 G for 10 minutes. The content of 1 centrifuge tube was divided into supernatant and sediment as in Comparative Example 35, and a didno concentration was measured about each of them. Adding 100 ml of delonized water, 1 g of sodium persulfate and 0.6 g of sodium hydroxide into the sediment in the other centrifuge tube, they were reacted keeping the temperature at 70 °C for 24 hours. The reaction was named as Example 54 and a dioxin concentration was measured about the reactant. The measured results are shown Table 3.

Table 20

40	Sample Condition	Category	Measured amounts of Dioxin (ng / pipe)	Decomposition ratio (%)
	Comparative Example 35	Supernatant	Under analytical sensitivity	•
45		Sediment	6. 1	-
	Example 54	Sediment after decomposition	0.4	93

## INDUSTRIAL APPLICABILITY

[0160] The present invention enables to provide a method of treating wastewater containing hardly decomposable harmful substances which separates the hardly decomposable harmful substances from wastewater by coagulation or adsorption and efficiently decomposes the separated hardly decomposable harmful substances in the solid condition without carrying out desorption operation.

#### Claims

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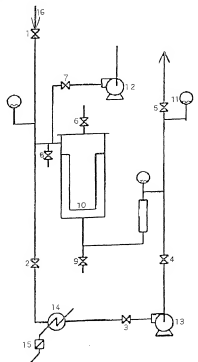
- A method of treating hardly decomposable harmful substance comprising:
- (A) a step for adding coagulant into wastewater containing a hardly decomposable harmful substance, (B) a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from a liquid, and
  - (C) a step for decomposing the hardly decomposable harmful substance by bringing peroxide into contact with the separated solid.
- The method of treatling hardly decomposable harmful substance according to Claim 1, wherein the hardly decomposable harmful substance is at least one kind selected from the group consisting of dibenzodoxin haildes, clibenzofuran haildes, polychlorinated biphenyls, benzene haildes, alklyphenols, phonol haildes, alkane haildes, alkene haildes, pithalic esters, bisphenols and polycyclic aromatic hydrocarbons.
- The method of treating hardly decomposable harmful substance according to Claim 1, wherein the coagulant employed in step (A) is at least one kind selected from the group consisting of polyacrylic acid, polyacrylamide resin, ferrous coagulant and aluminum-based coagulant.
- 4. The method of treating hardly decomposable harmful substance according to Claim 3, wherein the ferrous coagulant is at least one kind selected from the group consisting of ferric sulfate, polyferric sulfate, ferric chloride. and polyferric chloride.
- The method of treating hardly decomposable harmful substance according to Claim 3, wherein the aluminumbased coagulant is at least one of aluminum chloride and polyaluminum chloride.
  - The method of treating hardly decomposable harmful substance according to Claim 1, further adding coagulation promoter in the step (A).
- The method of treating hardly decomposable harmful substance according to Claim 6, wherein the coagulation
  promoter is an inorganic porous article or an organic porous article.
  - The method of treating hardly decomposable harmful substance according to Claim 6, wherein the coagulation promoter is at least one kind selected from the group consisting of ion exchange resin, diatom earth, activated clay, zeolite, pearlite and activated carbon.
  - The method of treating hardly decomposable harmful substance according to Claim 1, wherein a filtration method, a centrifugal separation method or a membrane separation method is employed in the step (B).
- 40 10. The method of treating hardly decomposable harmful substance according to Claim 1, wherein the peroxide used in the step (C) is an oxidizing agent.
  - 11. The method of treating hardly decomposable harmful substance according to Claim 10, wherein the oxidation agent is at least one of persulfate and permanganate.
  - 12. The method of treating hardly decomposable harmful substance according to Claim 11, wherein the persulfate is at least one kind selected from the group consisting of ammonium persulfate, sodium persulfate and potassium persulfate.
- 50 13. The method of treating hardly decomposable harmful substance according to Claim 11, wherein the permanganate is at least one of sodium permanganate and potassium permanganate.
  - 14. The method of treating hardly decomposable harmful substance according to Claim 1, wherein the peroxide used in the step (C) is at least one selected from the group consisting of salt of peroxo acid, hyperoxide and organic peroxide.
  - 15. The method of treating hardly decomposable harmful substance according to Claim 14, wherein the salt of peroxo acid is perborate or organic peroxide.

- 16. The method of treating hardly decomposable harmful substance according to Claim 14, wherein the hyperoxide is at least one selected from the group consisting of potassium hyperoxide, sodium hyperoxide, barlum hyperoxide and magnesium hyperoxide.
- 7 17. The method of treating hardly decomposable harmful substance according to Claim 14, wherein the organic peroxide is ditertiarybutylperoxide or dicumyl peroxide.
  - 18. A method of treating hardly decomposable harmful substance comprising:
- (a) a step for adding an adsorbent into wastewater containing a hardly decomposable harmful substance, (B) a step for performing solid-liquid separation which separates a solid containing the hardly decomposable harmful substance from all louid, and
  - (C) a step for decomposing the hardly decomposable harmful substance by bringing peroxide into contact with the separated solid.
  - 19. The method of treating hardly decomposable harmful substance according to Claim 18, wherein the adsorbent is at least one selected from the group consisting of ion exchange resin, diatom earth, activated day, zeolite, pearlite, titania and activated caton.
- 20 20. A method of treating a hardly decomposable substance wherein the hardly decomposable substance adsorbed to a solid is brought into contact with peroxides in an amount of 100 moles per the hardly decomposable substance without conducting any desorption operation.
- 21. The method of treating a hardly decomposable substance according to Claim 20, wherein the solid is any one of soll, sludge, incinerated ash or adsorbent.
  - 22. The method of treating hardly decomposable harmful substance according to Claim 20, wherein the hardly decomposable substance is at least one selected from the group consisting of dibenzolloxin halides, dibenzofutan halides, polychiofinated biphenyls, benzene halides, alkyphenols, phenol halides, alkane halides, alkane halides, alkane halides, alkane halides, polychiofinated biphenyls, benzene halides, alkane halides, alkane halides, alkane halides, polychiofinated biphenyls, benzenet halides, alkane halides, alkane halides, alkane halides, alkane halides, polychiofinated biphenyls, benzenet halides, alkane halides, alkane halides, alkane halides, alkane halides, polychiofinated biphenyls, benzenet halides, benzenet hali
  - 23. The method of treating hardly decomposable harmful substance according to Claim 20, wherein the peroxide is at least one selected from the group consisting of salt of peroxo acid, hyperoxide and organic peroxide.
- 24. The method of treating hardly decomposable harmful substance according to Claim 23, wherein the salt of peroxo acid is at least one selected from the group consisting of persulfate, permanganate, perborate and organic peroxide.
- 25. The method of treating hardly decomposable harmful substance according to Claim 24, wherein the persuifate is at least one selected from the group consisting of ammonium persuifate, sodium persuifate and potassium persuifate.
  - 26. The method of treating hardly decomposable harmful substance according to Claim 24, wherein the permanganate is any one of sodium permanganate and potassium permanganate
- 45 27. The method of treating hardly decomposable harmful substance according to Claim 23, wherein the hyperoxide is at least one selected from the group consisting of potassium hyperoxide, sodium hyperoxide, barium hyperoxide and magnesium hyperoxide.
- 28. The method of treating hardly decomposable harmful substance according to Claim 23, wherein the organic peroxide is ditertiarybutylperoxide or dicumyl peroxide.

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Fig. 1

## Waste water to be treated



## INTERNATIONAL SEARCH REPORT

International application No. PCT/JP03/09881

A. CLAS: Int.	SIFICATION OF SUBJECT MATTER CL <sup>7</sup> C02F1/52, 1/56, 1/28, B09	B3/00, A62D3/00, B01J20	/34					
According t	According to International Patent Classification (IPC) or to both national classification and IPC							
B. FIELD	S SEARCHED							
Minimum d Int.	locumentation searched (classification system followed C1' C02F1/52, 1/56, 1/28, B09	by classification symbols) B3/00, A62D3/00, B01J20,	/34					
Jits Koka:	Decumentation searched other than minimum documentation to the extent that such documents are included in the fields searched Jitsuyo Shinana Robo 1994-2003 Rokai Jitsuyo Shinana Robo 1994-2003 Rokai Jitsuyo Shinana Robo 1971-2003 Jitsuyo Shinana Toroku Robo 1970-2003 Electronic data base consulted during the international search (name of data base and, where praceicable, search terms used)							
C. DOCU	MENTS CONSIDERED TO BE RELEVANT							
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.					
X Y	JP 2002-239597 A (Hitachi Pl Construction Co.), 27 August, 2002 (27.08.02), Full text; Figs. 1 to 3 (Family: none)	Lant Engineering &	1-3,9,10, 20-22 4-8					
P,Y	JP 2003-033605 A (Miura Co., 04 February, 2003 (04.02.03) Full text (Family: none)	1-19						
P,Y	JP 2003-033772 A (Miura Co., 04 February, 2003 (04.02.03) Full text (Family: none)		1-19					
			-					
× Furth	er documents are listed in the continuation of Box C.	See patent family annex.						
"A" docume conside "E" carlier e date "L" docume cited to special docume means	contempts of Cliefs decomment:  In the distinging the powers instead of the set which is not one of the operational reflevance and to be of particular reflevance consument but published on or after the international filing net which may throw doubts on priority claim(s) or which is created the published on or after the custom or other reason (as specifical)  referring to no and disclosure, use, architishon or other reason (as specifical)  referring to no and disclosure, use, architishon or other reason published prior to the international filing dute but later reason (as published prior to the international filing dute but later	It is not accused published after the international filtring date or principly date and no in condicts with the expellentation but crited to make a final control of the expellentation but crited to considered not one of the considered not considered not not considered not con						
Date of the a	this to privrily date claimed.  Date of mainling of the international search 10 October, 2003 (10.10,03)  Date of mailing of the international search 28 October, 2003 (28.10.03)							
	ailing address of the ISAV nese Patent Office	Authorized officer						
Facsimile No	0.	Telephone No.						

Form PCT/ISA/210 (second sheet) (July 1998)

## INTERNATIONAL SEARCH REPORT

International application No. PCT/JP03/09881

ategory*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Y	JF 2002-320974 A (Ebara Corp.), 05 November, 2002 (05.11.02), Full text; Fig. 1 (Family: none)	1-19
Y	JF 2002-186963 A (Hitachi Zosen Corp.), 02 July, 2002 (02.07.02), Full text; Figs. 1 to 8 (Family; none)	1-19
¥	JP 2002-159975 A (Hitachi Zosen Corp.), 04 June, 2002 (04.05.02), Full text; Figs. 1 to 3 (Family: none)	1-19
Y	JP 2002-153891 A (Mitsubishi Heavy Industries, Ltd.), Ltd.), Ltd.), 28 May, 2002 (28.05.02), Full text: Figs. 1 to 4 (Family: none)	1-19
Y	JF 2002-153866 A (Kurita Water Industries Ltd.), 28 May, 2002 (28.05.02), Full text; Fig. 1 (Family: none)	1-19
Y	JP 2002-153703 A (Miura Co., Ltd.), 28 May, 2002 (28.05.02), Full text; Fig. 1 (Family: none)	1-19
Y	JP 2001-232356 A (Kurita Water Industries Ltd.), 28 August, 2001 (28.08.01), Full text, Figs. 1 to 3 (Family: none)	18,19
Y	JP 2001-149949 A (Shinko Pantec Co., Ltd.), 05 June, 2001 (05.06.01), Full text; Figs. 1 to 2 (Family: none)	1-19
Y	JP 2000-218278 A (Kurita Water Industries Ltd.), 08 August, 2000 (08.08.00), Full text (Family: none)	1-19
P,X P,Y	JP 2003-200134 A (Kurita Water Industries Ltd.), 15 July, 2003 (15.07.03), Full text (Family: none)	20-26 1-19,27,28

Form PCT/ISA/210 (continuation of second sheet) (July 1998)

## INTERNATIONAL SEARCH REPORT

International application No. PCT/JP03/09881

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X X	JP 2003-001277 A (Mitsui Engineering & Shipbuliding Co., Led.), 07 January, 2003 (07.01.03), Full text (Family: none)	20-23,27 1-19,24-26, 28
X	JP 2001-179211 A (Miura Co., Ltd.), 03 July, 2001 (03.07.01), Full text (Family: none)	20-23,27 1-19,24-26, 28
X Y	JP 2002-035772 A (Ueda Textile Science Foundation), 05 February, 2002 (05.02.02), Full text; Fig. 1 (Family: none)	20-24 1-19,25-28
Y	JF 2001-179269 A (Kurita Water Industries Ltd.), 03 July, 2001 (03.07.01), Full text (Family: none)	1-28
Х	JP 2002-355638 A (Mitsui Engineering & Shipbuilding Co., Itd.), 10 December, 2002 (10.12.02), Full text; Figs. 1 to 5 (Family: none)	20-23,27 24-26,28

Form PC1/ISA/210 (continuation of second sheet) (July 1998)

## INTERNATIONAL SEARCH REPORT

International application No. PCT/JP03/09881

BOX 1 Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)
This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.:
because they relate to subject matter not required to be searched by this Authority, namely:
÷
2. Claims Nos.:
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be curried out, specifically:
3. Claims Nos.:
because they are dependent claims and are not drafted in accurdance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 3 of first sheet)
The international Searching Authority found multiple inventions in his international application, as follows: The inventions claimed in claims 1-17, 18, 19 and 20-28 are common in having the technical feature of bringing a solid into contact with a peroxide to thereby decompose hardly decomposable harmful substances.  Nowever, this technical feature is known as described in JP 2001-179211 a Therefore, among this group of inventions, there is no technical relationship expressed by the same or corresponding special technical features. Consequently, it cannot be recognized that the above group of inventions are so linked as to form a single general inventive concept.
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
X as all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
<ol> <li>As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos:</li> </ol>
No required additional search fees were finely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the cidines; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protes: accompanied the payment of additional search fees.
X

Form PCT/ISA/210 (continuation of first sheet (1)) (July 1998)